Temporal dynamics of natural communities of marine algal viruses and eukaryotes

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ABSTRACT: The composition of algal virus communities in relation to temperature, salinity, chlorophyll a (chl a) concentration and eukaryotic community composition was monitored at a single location for 14 mo. Changes in algal virus and eukaryote communities were determined using polymerase chain reaction (PCR) and denaturing gradient gel electrophoresis (DGGE) to generate genetic fingerprints. Sequence analysis of bands extracted from denaturing gradient gels revealed the presence of at least 5 distinct viruses as well as temporally dynamic and diverse communities of eukaryotes that included taxa from the viridiplantae, fungi and metazoa. Comparison of algal virus fingerprints with environmental conditions revealed that, at certain times, changes in algal virus community composition were coincident with changes in tide height, salinity or chl a concentration. However, algal virus community changes were not often coupled to eukaryote community changes. The lack of coincidence between changes in virus and eukaryote communities can be explained by the presence of organisms that were not hosts of the detected viruses. It is likely that the uncoupling of 18S and AVS fingerprints was due to succession among non-host eukaryotes. Although algal virus fingerprint patterns were stable throughout most of the study, stable eukaryote fingerprint patterns were observed only during the winter months. Furthermore, specific taxa of algal viruses persisted in fluctuating physical and biological environments. We concluded that the constant production of, and mortality from, some taxa of algal viruses provide further evidence that algal viruses affect phytoplankton community structure and dynamics.

KEY WORDS: Algal viruses · Phycodnaviridae · Eukaryotes · DGGE · Molecular fingerprints · Diversity

INTRODUCTION

After the discovery that viruses were abundant in the ocean, marine microbiologists quickly recognized their ecological importance (Thingstad et al. 1993, Bratbak et al. 1994, Suttle 1994). There is also mounting evidence that viruses have a significant effect on oceanic energy and material flow (Fuhrman 1999, Wilhelm & Suttle 1999). Moreover, there is now compelling data showing that viruses are central players affecting phytoplankton communities in the sea (Suttle 2000). An important contribution to the study of viruses that infect marine phytoplankton was the development of polymerase chain reaction (PCR) techniques that are specific for algal viruses belonging to the Phycodnaviridae (Chen & Suttle 1995). Using these culture-independent PCR methods, sequences belonging to unknown taxa of algal viruses were amplified from the Gulf of Mexico demonstrating the utility of this molecular approach (Chen & Suttle 1996). Subsequently, we developed a denaturing gradient gel electrophoresis (DGGE) protocol to resolve sequences amplified with the algal-virus-specific (AVS) PCR primers (Short & Suttle 2000). Recently, molecular fingerprinting of algal virus communities facilitated the recovery of algal virus gene sequences from different environments revealing that groups of closely related algal viruses were geographically widespread (Short & Suttle 2002). The results of that study demonstrated that although further refinement of the PCR and DGGE methods was necessary, these methods could be applied to the study of spatial changes in algal virus community composition.

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The diversity of complex microbial populations was first studied using DGGE when Muyzer and co-workers obtained molecular fingerprints of 16S rDNA fragments amplified from an aerobically grown bacterial biofilm (Muyzer et al. 1993). Since then, DGGE has become a common method to obtain molecular fingerprints and has been used to study microbial diversity in many different contexts. For the most part, DGGE has been applied to the study of bacterial community composition as determined by PCR amplification of 16S rDNA. DGGE has been used to examine the microbial community diversity of biostimulated shorelines (Ogino et al. 2001), hot spring microbial mats (Ferris et al. 1997), California estuaries (Murray et al. 1996) and the Arctic Ocean (Ferrari & Holibaugh 1999). In addition, DGGE has been applied to examine diversity in marine cyanophage (Wilson et al. 2000) and algal virus communities (Short & Suttle 1999, 2002). Recently, PCR and DGGE fingerprinting methods have also been developed to examine the diversity of eukaryotic microbes (Van Hannen et al. 1998, Diz et al. 2001). These eukaryote fingerprinting techniques have been applied to investigate the temporal variability of marine eukaryotes in an enclosure (Castberg et al. 2001) and the spatial variability through a salinity gradient in a solar saltern (Casamayor et al. 2002). However, to our knowledge, molecular fingerprinting has not been used to examine temporal changes in marine eukaryotic communities.

The primary goal of this study was to determine the annual variability in the genetic fingerprints of natural algal virus communities over a 64 wk period. Indirect evidence of changes in algal virus community composition has been noted when blooms of the alga *Emiliania huxleyi* have been succeeded by an increase in large virus-like particles or *E. huxleyi* viruses (Bratbak et al. 1993, Castberg et al. 2001). Furthermore, changes in aquatic viral assemblages have been examined by comparing genome size classes with pulsed field gel electrophoresis (Wommack et al. 1999, Steward et al. 2000). However, to date, no direct examination of temporal changes in algal virus community composition has been conducted. A secondary goal of this study was to relate changes in algal virus diversity, as inferred from molecular fingerprints, to changes in eukaryote diversity and the physical environment. The results of this study show that although the temporal changes of algal viruses and eukaryotes were not closely coupled, changes were evident in both communities. More importantly, this study demonstrates that changes in the composition of marine algal virus communities often coincide with changes in the environment.

**MATERIALS AND METHODS**

**Sample collection.** Using a bucket and funnel, 40 l samples were collected weekly from the surface water at Jericho Pier, Vancouver, Canada (49° 11’ N, 123° 10’ W). Sample temperature and salinity were measured approximately 15 min after collection, and duplicate acetone extracted chlorophyll *a* (chl *a*) samples were measured fluorometrically using established protocols (Parsons et al. 1984). The virus-size fraction was concentrated using previously described methods (Suttle et al. 1991, Short & Suttle 2002). Briefly, samples were pressure-filtered (<17 kPa) through 142 mm in diameter glass fiber (GC50; nominal pore size 1.2 µm, Advantec MFS) and PVDF filters (GVWP; 0.45 µm pore size, Millipore) connected in series. Remaining particulate material in the filtrates was concentrated to a final volume of ca. 200 ml using a 30 kD cutoff, Amicon S1Y30 (Millipore) cartridge. Concentrated samples were stored at 4°C in the dark. The glass fiber and PVDF filters were stored at –20°C in the dark until nucleic acid extraction.

**DNA extraction and PCR.** Nucleic acids were extracted from virus-size-fraction concentrates, glass-fiber filters and PVDF filters. Thirty-five ml subsamples of the virus-size concentrates were centrifuged in an RC80 ultracentrifuge with an AH629 rotor (Sorvall®) at 28 500 rpm for 3.5 h at 20°C. Nucleic acids were extracted from the resuspended pellets (100 µl) using an established hot/cold technique (Chen et al. 1996). DNA was also extracted from aseptically excised 2 cm² portions of the 142 mm in diameter filters. After Díez et al. (2001), filter subsamples were incubated at 37°C for 45 min in 500 µl lysozyme (final concentration: 1 mg ml⁻¹) and at 65°C for 1 h after a 500 µl solution of sodium dodecyl sulfate (final concentration: 1%) and Proteinase K (final concentration: 0.2 mg ml⁻¹) was added. DNA was purified from the filter lysates by sequential extractions with equal volumes of phenol, phenol-chloroform-isooamyly alcohol (25:24:1) and chloroform-isoamyly alcohol (24:1). Purified DNA was precipitated and washed with ethanol as described (Sambrook et al. 1989). Extracted DNA was stored at –20°C in 10 mM Tris-HCl (pH 8.5) until further use.

PCR annealing temperature optimization was necessary to improve the specificity of algal-virus polymerase amplification. From the sample collected on February 1, 2001, 3 µl (ca. 0.5% of the 40 l sample) of virus-size extract was added to a 47 µl PCR mixture containing *Tag* DNA polymerase assay buffer (50 mM KCl, 20 mM Tris-HCl, pH 8.4), 1.5 mM MgCl₂, 0.16 mM of each deoxyribonucleoside triphosphate (dNTP), 10 pmol of the primer AVS1 and 30 pmol of AVS2, and 0.625 U of PLATINUM® *Tag* DNA polymerase (Invitrogen Life Technologies). The degenerate algal-virus-specific primers AVS1 and AVS2 amplify a 700 bp frag-
ment of pol (Chen & Suttle 1995). The negative control contained all reagents except template and was placed at the lowest annealing temperature. PCR was carried out in a Hybaid PCR Express thermal cycler with the following cycle parameters: denaturation at 95°C for 90 s; 35 cycles of denaturation at 95°C for 45 s; annealing gradient from 42 to 56°C for 45 s; and extension at 72°C for 45 s with a final 5 min extension.

Once the PCR annealing temperature was optimized, algal virus pol was amplified using 2 separate PCR reactions. First, 3 µl of virus-size extract was added to PCR mixtures as described above. All reaction conditions were as above except the annealing temperature was 45°C and only 32 cycles were performed. After electrophoresis, a disposable Pasteur pipette was used to excise a plug from the 700 bp band amplified in each reaction. The plug was placed in 200 µl of sterile 1.0 × TAE (40 mM Tris-base, 20 mM sodium acetate, 1 mM EDTA, pH 8.5) and heated to 65°C for 20 min to elute DNA. In the second-stage PCR, 2 µl of the eluant was used to increase product yields for the subsequent DGGE analysis. These reactions were identical to first-stage PCR except the number of cycles was limited to 20. Second-stage negative controls were conducted using the eluant of plugs excised from the negative control lane of gels of first-stage PCR.

Eukaryotic 18S DNA fragments were amplified from filter extracts using previously described universal eukaryotic primers Euk1A, Euk516r and Euk516r-GC (Diez et al. 2001). Five µl of a filter extract were added to a 45 µl PCR mixture containing Taq DNA polymerase assay buffer (50 mM KCl, 20 mM Tris-HCl, pH 8.4), 1.5 mM MgCl₂, 0.16 mM of each deoxyribonucleoside triphosphate (dNTP), 15 pmol of the 18S primers Euk1A and Euk516r, and 0.625 U of PLATINUM® Taq DNA polymerase. PCR was carried out using a Hybaid PCR Express thermal cycler with the following cycle parameters: denaturation at 95°C for 90 s; 32 cycles of denaturation at 95°C for 1 min; annealing at 56°C for 1 min; and extension at 72°C for 1 min with a final 5 min extension. Like the algal virus pol reactions, a second PCR reaction was conducted using first-stage gel plugs as templates. In this case, second-stage reactions were used to incorporate a GC clamp onto the PCR products for the subsequent DGGE analysis. All second-stage PCR conditions were the same as first-stage 18S amplification except the downstream primer Euk516r-GC was used and the reaction was limited to 20 cycles.

PCR products were electrophoresed in 1.8% LE agarose (FMC BioProducts) in 1.0 × TAE at 90 V for 75 min. Gels were stained with ethidium bromide and photographed on an ultraviolet transilluminator with a Nikon Coolpix 950 digital camera. Digital images were inverted and contrast and brightness was adjusted using Adobe Photoshop 5.0 LE.

**DGGE and sequencing.** Initially, DGGE conditions were optimized for maximum resolution by electrophoresing several samples in different gradient gels. From these gels, bands were excised, eluted and amplified as described above for the templates of second-stage PCR. For each type of DGGE analysis (AVS or 18S), 6 bands were selected and pooled for use as standards in subsequent gradient gels. DGGE of second-stage AVS PCR products was conducted using 20 to 40% denaturing gradients cast in 7 to 8% gradient polyacrylamide gels. Approximately 50 µl (range 35 to 60) of the pooled products of 3 separate second-stage reactions were loaded into wells with 5 µl of 10.0 × loading buffer (50% v/v glycerol, 0.33 M EDTA, pH 8.0, and 0.08% w/v bromphenol blue). The amount of pooled PCR products loaded into each lane was normalized after the products were quantified in agarose gels using the program Gel-Pro® Analyzer (Media Cybernetics). Wells flanking the samples were loaded with 45 µl of pooled AVS standards. Electrophoresis was carried out for 15 h in 1.0 × TAE at 80 V and a constant temperature of 60°C using the D-code™ electrophoresis system (Bio-Rad Laboratories). Similarly, DGGE of the second-stage 18S PCR products was conducted using 25 to 47.5% denaturing gradients cast in 6% polyacrylamide gels. From 40 to 50 µl of pooled PCR products of 3 separate second-stage reactions was loaded into each well with 10 × loading buffer; the amount of DNA loaded was normalized as described for AVS-DGGE. Wells flanking the samples were loaded with 40 µl of pooled 18S standards. Electrophoresis was carried out for 15 h in 1.0 × TAE at 100 V and 59°C. Gels were stained in a 0.1 × SYBR® Green I (Molecular Probes) solution overnight, de-stained in water for at least 30 min and photographed as described above for agarose gels. Digital images of the denaturing gradient gels were analyzed using the program GelCompar II (Applied-Maths BVBA). For each type of DGGE analysis (AVS or 18S), band positions in individual gel images (6 AVS and 12 18S) were normalized using lanes loaded with the standards. To estimate sample similarities within each fingerprint set (AVS, 18S 0.45 to 1.2 µm fraction, and 18S > 1.2 µm fraction) Pearson product moment correlation coefficients were calculated from lane densitometry plots and were used to calculate UPGMA dendrograms.

Bands from different gel positions were excised from the AVS or 18S gradient gels for sequencing. Gel slices were then placed in sterile microcentrifuge tubes with 200 µl of 1.0 × TAE, heated to 95°C for 5 min to elute the DNA, and stored at –20°C until further use. Eluted DNA fragments from 10 arbitrarily selected AVS bands were amplified as described above for second-stage AVS reactions. Amplified AVS DNA was then cloned using a pGEM®-T Easy kit (Promega). Eluted DNA...
fragments from 19 arbitrarily selected 18S bands were amplified as described for second-stage 18S reactions. AVS plasmid DNA (300 to 500 ng) or 18S amplified DNA (4 µl) and an appropriate oligonucleotide primer (M13 forward/reverse or Euk1A/S16r, respectively) were added to sequencing reactions using AmpliTaq FS® BIGDYE Terminator cycle sequencing chemistry (Applied Biosystems) and excess dye terminators were removed using CENTRI-SEP™ spin columns (Princeton Separations). Reactions were run in ABI Model 373 Stretch or ABI Prism 377 automated sequencers (Applied Biosystems).

Algal-virus pol sequences obtained from DGGE bands were compared to known virus pol genes available in GenBank (accession numbers listed below). Sequences were edited using BioEdit version 5.0.7 (Hall 1999) and inferred amino acids were aligned using the sequence alignment program CLUSTAL X (Thompson et al. 1997) with default settings. The alignment of the resulting 340 amino acid positions was used to construct a maximum likelihood tree using the program TREE-PUZZLE version 5.0 (Strimmer & von Haeseler 1996) and phylogenetic trees were drawn using the program TreeView (Win32) version 1.6.1 (Page 1996). The sequences recovered from the 18S gels were compared to nucleotide sequences in GenBank (available at www.ncbi.nlm.nih.gov) using blastn and small subunit rDNA sequences in the Ribosomal Database Project II (available at rdp.cme.msu.edu/html) using Sequence Match (Maidak et al. 2001).


The accession numbers of virus pol sequences used in the phylogenetic analysis are: ASFV, X73330; BSA99-1, AF405577; BSA99-2, AF405578; BSA99-5, AF405581; BSB99-2, AF405588; ChV-PW1, U32983; ChV-PW3, U32984; CVA-1, U32985; EBV, V01555; EsV-1-93, AAK14511; FSV, AAB67116; HSV-1, X04771; HCMV, M14709; MVB99-2, AF405592; MvP-GM1, U32977; MvP-P88, U32980; MpV-PL1, U32982; MpV-SG1, U32981; MpV-S61, U32975; NY-2A, M86837; OTU1, U36931; OTU2, U36932; OTU3, U36933; OTU4, U36934; OTU5, U36935; PBCV-1, M86836; PSB99-1, AF405593; PSC99-1, AF405597; PSC99-2, AF405598; SIA99-1, AF405590; SO98-1, AF405572; SO98-2, AF405573; SO98-3, AF405574; SO98-5, AF405576.

**RESULTS**

Optimization of algal-virus pol PCR successfully eliminated a previously observed 550 bp artifact (Short & Suttle 2002). At annealing temperatures above 48°C, PCR reactions with the February 1, 2001 sample and AVS primers produced 700 and 550 bp bands. The highest yield of the 700 bp band was at the lowest annealing temperature and diminished with increasing temperature, while the highest yield of the 550 bp band was at the highest annealing temperature and disappeared as the temperature decreased to 46°C (Fig. 1). Two-stage AVS PCR using 45°C annealing temperatures produced only bright 700 bp bands. Similarly, second-stage PCR with 18S primers produced only the expected 550 bp DNA fragments (data not shown).

By sequencing and phylogenetic analysis, 10 bands from the AVS fingerprints were characterized. All 10 sequences recovered from the AVS fingerprints encoded the purported pol motif YGDTDS, and comparison to other virus pol sequences revealed that their inferred amino acid sequences fell within the clade of known algal viruses belonging to the Phycodnaviridae (Fig. 2). Although none of the Jericho Pier sequences were identical, the percentage of identical nucleotides was at least 98 among the sequences JPavs-37, JPavs-42, JPavs-51 and JPavs-52. Similarly, the sequences JPavs-64 and JPavs-65 were 99.5%, and JPavs-22 and JPavs-38 were 98.0% identical. The remaining sequences (JPavs-53 and JPavs-55) were less than 89% identical to any other sequence. Several sequences were more than 98% identical to other environmental sequences available in GenBank. For example, the
sequences JPsavs-64 and JPsavs-65 were at least 99% identical to the sequence BSA99-1, and the sequence JPsavs-53 was 99% identical to SO98-5. Only 1 sequence from the AVS fingerprints (JPsavs-55) was less than 90% identical to any known algal virus pol fragment. Nineteen bands excised from the 18S fingerprints were sequenced and compared to sequence databases. When compared to nucleotide sequences in GenBank, all of the sequences recovered from the 18S gradient gels were related to 18S rDNA sequences. The Jericho Pier 18S sequences were related to 18S rDNA sequences from several eukaryote lineages including Alveolata, Fungi, Metazoa, Stramenopiles and Viridiplantae (Table 1).

The salinity and temperature of the water at Jericho Pier varied seasonally with the highest temperature and lowest salinity occurring during summer, and the opposite is true of the winter months (Fig. 3). During winter, temperature and salinity fluctuated less than at other times of the year. With the exception of the winter months, when it was lowest, chl a concentrations were variable. Because the local tidal cycle is mixed diurnal, the tide height at Jericho Pier varied considerably among the times of sample collection.

Visual inspection of the composite DGGE images revealed that the banding patterns of the AVS fingerprints (Fig. 3) appeared less variable than the 18S fingerprints (Fig. 4). Throughout most of the study period, especially during the spring, the 18S fingerprint patterns changed from week to week while the AVS patterns were relatively constant. Nonetheless, bands present throughout most of the sample period were observed in all 3 fingerprints. The bands indicated at Position 1 in the AVS fingerprint can be observed in all but a few samples (Fig. 3), and Bands euk51 and euk113 were present in the majority of the 18S samples from the 0.45 to 1.2 and >1.2 µm fractions respectively (Fig. 4). On the other hand, some bands were observed in only a few samples. Band avs55 was observed only during the winter and early spring of each year; this band was present from the beginning of the study until April 7, 2000 and reappeared from January 25, 2001 until April 4, 2001. In addition, Band avs53 was present only in February and March 2001, and Bands 1 and 2 in the 0.45 to 1.2 and >1.2 µm 18S fractions (Fig. 4) appeared only once. Overall, patterns of band appearance and disappearance were observed in all fingerprints.

Visual inspection also revealed changing fingerprint patterns coincident with changes in the physical characteristics of samples from Jericho Pier. From November 2000 through February 2001, the salinity was >25 ppt, the temperature was <15°C and chl a concentrations were <3 µg l⁻¹. During these months, the 18S fingerprints of
Table 1. Results of 18S ribosomal database searches with Jericho Pier sequences

<table>
<thead>
<tr>
<th>Sequence</th>
<th>Fraction</th>
<th>NCBI blastn results$^a$</th>
<th>RDP II sequence match results$^b$</th>
</tr>
</thead>
<tbody>
<tr>
<td>JPeuk-1</td>
<td>0.45–1.2 µm</td>
<td>Eukaryota, Stramenopiles, Uncultured Stramenopile BAQA21, AF372755 (771)</td>
<td>Eukaryota, Stramenopiles, Uncultured Stramenopile, OLI61105, AF167414 (0.642)</td>
</tr>
<tr>
<td>JPeuk-2</td>
<td>0.45–1.2 µm</td>
<td>Alveolata, Ciliophora, Spirotrichea, Stylophaga mytilus, AJ310499 (622)</td>
<td>Alveolata, Ciliophora, Spirotrichea, Stylophaga mytilus, AJ310499 (0.588)</td>
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<tr>
<td>JPeuk-4</td>
<td>0.45–1.2 µm</td>
<td>Fungi, Ascomycota, mitosporic Ascomycota, Geosmithia viridis, AB033527 (991)</td>
<td>Fungi, Ascomycota, Pezizomycotina, Talaromyces flavus, M83262 (0.901)</td>
</tr>
<tr>
<td>JPeuk-6</td>
<td>0.45–1.2 µm</td>
<td>Viridiplanta, Streptophyta, Embryophyta, Casuarina equisetifolia, U42515 (983)</td>
<td>Viridiplanta, Streptophyta, Embryophyta Fagus grandifolia, AF206910 (0.871)</td>
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<tr>
<td>JPeuk-20</td>
<td>0.45–1.2 µm</td>
<td>Viridiplanta, Chlorophyta, Prasinophyceae, Prasinophyta symbiont of radiolarian, AF166380 (922)</td>
<td>Prasinophyta radiolarian symbiont, AF166379 (0.783)</td>
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<td>JPeuk-27</td>
<td>0.45–1.2 µm</td>
<td>Metazoa, Arthropoda, Crustacea, Neocalanus tansus, X85393 (833)</td>
<td>Metazoa, Brachiopoda, Linguliformea, Lingula anatine, AB033527 (0.868)</td>
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<td>JPeuk-38</td>
<td>0.45–1.2 µm</td>
<td>Viridiplanta, Chlorophyta, Prasinophyceae, Mantoniella Antarctica, AB017128 (795)</td>
<td>Viridiplanta, Chlorophyta, Prasinophyceae Ostreococcus tauri, Y15814 (0.608)</td>
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<td>JPeuk-40</td>
<td>0.45–1.2 µm</td>
<td>Eukaryota, environmental samples, Eukaryote marine clone MEI-1, AF363153 (932)</td>
<td>Eukaryota, environmental samples, Lobosphaera tiroleis, AB000651 (0.713)</td>
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<td>JPeuk-51</td>
<td>0.45–1.2 µm</td>
<td>Eukaryota, environmental samples, Eukaryote marine clone MEI-1, AF363153 (957)</td>
<td>Eukaryota, environmental samples, Lobosphaera tiroleis, AB000651 (0.716)</td>
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<tr>
<td>JPeuk-119</td>
<td>0.45–1.2 µm</td>
<td>Eukaryota, environmental samples, Eukaryote marine clone MEI-1, AF363155 (660)</td>
<td>Eukaryota, environmental samples, Coelastrella multistriata, AB012846 (0.487)</td>
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<td>JPeuk-59</td>
<td>&gt;1.2 µm</td>
<td>Stramenopiles, Bacillariophyta, Coscinodiscophyceae, Skeletonema pseudocostatum, X58393 (833)</td>
<td>Stramenopiles, Bacillariophyta, CoscinodiscophyceaeSkeletonema costatum, M54988 (0.761)</td>
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<td>JPeuk-65</td>
<td>&gt;1.2 µm</td>
<td>Eukaryota, environmental samples, Uncultured eukaryote clone LEMD134, AF372806 (321)</td>
<td>Eukaryota, environmental samples, Theileria sp., AB012202 (0.453)</td>
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<td>JPeuk-71</td>
<td>&gt;1.2 µm</td>
<td>Alveolata, Dinophyceae, Gymnodiniales, Amphidinium semilunatum, AF274256 (900)</td>
<td>Alveolata, Dinophyceae, Gymnodiniales Karlodinium galatheaen, KT-77B, AF172712 (0.738)</td>
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<td>JPeuk-77</td>
<td>&gt;1.2 µm</td>
<td>Viridiplanta, Chlorophyta, Trebouxiophyceae, Nannochloris sp. RICC 011, AJ131691 (977)</td>
<td>Viridiplanta, Chlorophyta, Trebouxiophyceae Nannochloris sp., RICC 011, AJ131691 (0.917)</td>
</tr>
<tr>
<td>JPeuk-80</td>
<td>&gt;1.2 µm</td>
<td>Eukaryota, environmental samples, Eukaryote clone OLI1151, AJ402343 (1007)</td>
<td>Eukaryota, environmental samples, Prorocentrum micans, OLI11511, AJ402343 (977)</td>
</tr>
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<td>JPeuk-84</td>
<td>&gt;1.2 µm</td>
<td>Alveolata, Ciliophora, Spirotrichea, Stylophaga mytilus, AJ310498 (400)</td>
<td>Alveolata, Ciliophora, Spirotrichea Haliella grandinella, AJ164137 (0.448)</td>
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<td>JPeuk-94</td>
<td>&gt;1.2 µm</td>
<td>Alveolata, Dinophyceae, Gymnodiniales, Amphidinium semilunatum, AF274256 (874)</td>
<td>Alveolata, Dinophyceae, Gymnodiniales Karlodinium galatheaen, AF172712 (0.756)</td>
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<td>JPeuk-108</td>
<td>&gt;1.2 µm</td>
<td>Fungi, Ascomycota, Pneumocystidomycetes, Nannochloris sp., RICC 011, AJ131691 (977)</td>
<td>Unidentified eukaryote 18S rRNA, AJ130849 (0.638)</td>
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<tr>
<td>JPeuk-113</td>
<td>&gt;1.2 µm</td>
<td>Eukaryota, environmental samples, Eukaryote clone OLI1151, AJ402343 (1013)</td>
<td>Eukaryota, environmental samples, Prorocentrum micans, M14649 (0.571)</td>
</tr>
</tbody>
</table>

$^a$Taxonomy, name, GenBank accession # and bit score of the highest scoring sequence is shown

$^b$Taxonomy, name, GenBank accession # and similarity value of the most similar sequence is shown

both fractions appeared less variable than at other times of the year (Fig. 4). Similarly, the AVS fingerprints remained relatively constant through the winter months, albeit with later timing than the 18S fingerprints; the AVS fingerprints changed little from the months, albeit with later timing than the 18S fingerprints (Lanes A and C, respectively, Fig. 3). Cluster analysis of each fingerprint set and inspection of the resulting UPGMA trees revealed that the AVS fingerprints were more similar to one another than were the 18S fingerprints (Fig. 5). With the exception of the samples collected on March 10 and May 19, 2000, samples from March to May 2000 (white bars, Fig. 5) clustered beside one another in the AVS tree, yet were distributed among 5 separate clusters in each 18S dendrogram. Similarly, the fingerprints from mid-
Fig. 3. Physical data and algal-virus-specific (AVS) fingerprints from Jericho Pier. Top: salinity, temperature and chlorophyll $a$ in weekly samples collected at Jericho Pier. Middle: composite image of AVS fingerprints. Annotations on the gel image show the positions of bands sequenced and lanes or bands discussed in the text. Bottom: predicted tidal heights at the time of sample collection are shown in the bottom panel. Sample dates are written as yy/mm/dd.
Fig. 4. Physical data and 18S rDNA fingerprints from Jericho Pier. Top: salinity, temperature and chlorophyll a in weekly samples collected at Jericho Pier. Middle: DGGE fingerprints of 18S rDNA amplified from 0.45 to 1.2 µm size fraction extracts. Bottom: DGGE fingerprints of 18S fingerprints from the >1.2 µm size fraction extracts. Annotations show the positions of bands sequenced and lanes or bands discussed in the text. Sample dates are written as yy/mm/dd.
Fig. 5. DGGE fingerprint cluster analysis. UPGMA trees of sample similarities for (A) the algal-virus-specific (AVS), (B) 0.45 to 1.2 µm 18S and (C) >1.2 µm 18S fingerprints. The scales to the right of each dendrogram show percent similarity. The resolution of each tree (maximum similarity) was estimated from the minimum similarity of standard lanes (data not shown). The black, grey and white bars below the dates of each dendrogram show the positions of sequences discussed in the text. Sample dates are written as yy/mm/dd.
September to early December (black bars, Fig. 5) and from February through March 2001 (grey bars, Fig. 5) formed a fewer clusters in the AVS dendrogram compared to 18S. In general, UPGMA analysis of pairwise densitometry correlations illustrated that AVS fingerprints were more similar to one another and samples from the same time of year tended to form fewer and more similar clusters when compared to 18S fingerprints.

**DISCUSSION**

The goal of this study was to observe changes in algal virus communities at Jericho Pier over several months and relate them to eukaryote diversity, chl a concentration, temperature, salinity and tidal height. Sequencing efforts revealed at least 5 distinct groups of viruses and several kingdoms of eukaryotes. We discovered that, at certain times, tide height and salinity changes were coincident with changes in the algal virus community composition. On 1 occasion, a phytoplankton bloom was also coincident with a shift in the algal virus community structure. In general, the composition of the detected algal virus communities was temporally less variable than that of the eukaryote communities.

Before algal virus communities could be monitored using molecular techniques, the fidelity of PCR with the primers AVS1 and AVS2 had to be improved. Established PCR conditions permitted amplification of sequences not related to algal virus pol fragments. Previously, we demonstrated that production of these PCR artifacts was due to amplification from the upstream primer AVS1 alone (Short & Suttle 2002). Because the melting temperature of AVS1 is ca. 10°C higher than AVS2, efforts to increase reaction specificity by increasing annealing temperature failed. Therefore, it is likely that increased reaction specificity at lower annealing temperatures was due to an increase in AVS2 annealing efficiency. It is notable that every band sequenced in this study fell within a monophyletic group of known algal virus pol sequences (Fig. 2), providing strong evidence that the fingerprints represented algal virus pol fragments only.

Many algal virus sequences recovered from Jericho Pier were more than 98% identical to each other and to sequences from other environmental samples. Two sets of sequences (JPavs64 & 65 and JPavs51 & 52) differed by only 2 and 3 nt, respectively. These sequences were recovered from bands at different positions in the fingerprint, and differences were due to nucleotides incorporated at sites of primer degeneracy. Therefore, quantitative examination of algal virus richness was not possible with degenerate primer PCR and DGGE.

Nonetheless, when primer sequences were excluded, no algal virus sequences were more than 99.8% identical, and it is unlikely that observed differences were due to PCR error alone. Using reported Taq error rates (Bracho et al. 1998), 1 misincorporated base per sequence can be estimated during algal virus pol amplification resulting in an identity difference of only 0.1%. Currently, we do not know if pol sequences differing by less than 1% represent different strains or ecotypes of viruses. Nonetheless, the 10 sequences recovered from Jericho Pier resolved onto 5 separate branches of the maximum likelihood tree (Fig. 2), and the minimum difference between any 2 of these branches was ca. 8%. Therefore, it is likely that each of the 5 branches represent distinct taxa of uncultivated viruses. All sequences recovered from the 18S fingerprints were related to small subunit rDNA sequences. While the majority of sequences recovered from the 18S fingerprints were related to phytoplankton, 3 sequences were related to zooplankton, 2 to fungi and 1 to a terrestrial plant. However, because only a limited number of bands were sequenced, the recovered sequences did not quantitatively represent the organisms present. Nonetheless, sequences from the 18S fingerprints demonstrated that several kingdoms of eukaryotes were detected with the described PCR methods.

During this study, changes in the virus fingerprints were often related to the physical characteristics of the water at Jericho Pier. The relationship between algal virus community dynamics and the physical environment was apparent when AVS fingerprints were compared to tide height. At certain times, changes in the tide height coincided with changes in the AVS fingerprint patterns. The bands indicated at Position 2 in the AVS fingerprints of samples collected from June 29 to August 10, 2000, alternated position coincident with changes in tide height (Fig. 3). However, at other times, fingerprint changes were not coincident with tide height changes. Intra-annual variability of the tidal influence on virus communities may be related to other aspects of the physical characteristics of the sample site. During the weeks that tide height changes were coincident with changes in the pattern of successive AVS fingerprints, the salinity was lowest indicating that the Fraser River freshet had peaked; the surface water at Jericho Pier is strongly influenced by freshwater input from the Fraser River (Harrison et al. 1983). During the freshet, freshwater surface flow may have caused more pronounced shifts in phytoplankton and virus community composition as the water at Jericho Pier ebbed and flooded.

Another example of an environmental influence on AVS fingerprint patterns was the change in the pattern coincident with 1 of the 3 phytoplankton blooms
observed during the study; during each bloom chl a concentrations exceeded 20 µg l⁻¹. These blooms occurred in summer 2000, autumn 2000 and early spring 2001 (Lanes A, B and C, Fig. 3). Examination of Lugol’s iodine-preserved samples (Parsons et al. 1984) revealed that the summer bloom was a nearly monospecific bloom of the Raphidophyte Heterosigma akashiwo. In contrast, the fall bloom was mixed and dominated by the diatoms Ditylum, Leptocylindrus and Pseudonitzschia, and the dinoflagellates Prorocentrum, Dinophysis and Protoperidinium; the 2001 spring bloom was dominated by the diatoms Skeletonema, Chaetoceros and Thalassiosira. The fact that the AVS fingerprint did not change during the summer Heterosigma bloom or the spring diatom bloom was explicable. Although isolates of viruses that infect Heterosigma are established in culture (Nagasaki & Yamaguchi 1997, Lawrence et al. 2001), we have not successfully amplified pol sequences from these viruses, suggesting that the AVS primers are not suitable. Also, to our knowledge, no viruses that infect diatoms have been isolated. During the mixed diatom and dinoflagellate bloom of the fall, the >1.2 µm fraction 18S fingerprint contained many bands, suggesting that the bloom was of a diverse group of phytoplankton (Lane A, Fig. 4). At the same time, the band pattern of the AVS fingerprints changed relative to the fingerprints from surrounding weeks. While this does not establish a causal relationship between changes in eukaryote richness and AVS fingerprints, the coincidence is intriguing; the presence of Phycodnaviridae hosts is most probable when eukaryote diversity was high. Because not all phytoplankton blooms were coincident with AVS fingerprint changes, it is likely that the viruses detected with the AVS primers do not infect the dominant bloom-forming algae at Jericho Pier.

In general, algal virus fingerprints were less variable than eukaryote fingerprints. This contrasting variability of the 2 communities was not surprising; the viruses detected with the AVS primers all belonged to the Phycodnaviridae (dsDNA viruses that infect eukaryotic phytoplankton), and therefore infected only a small subset of eukaryotes. Although the majority of sequences from the 18S fingerprints were related to phytoplankton, several were not. Because only a limited number of bands were sequenced from the 18S fingerprints, the ratios of taxa represented in the sequence data were not used to estimate the abundance of different eukaryote groups. Nonetheless, 18S sequences revealed the presence of many organisms that were not hosts of phycodnaviruses. The presence of non-host organisms may explain the lack of correspondence between virus and eukaryote community dynamics. We can speculate that changes in algal virus communities were coupled to changes in host communities. On the other hand, it is likely that changes in the 18S fingerprints due to the succession of non-host eukaryotes were not coupled to changes in the AVS fingerprints. In the future, development of PCR methods to fingerprint specific groups of phytoplankton, when combined with the virus fingerprinting methods outlined here, will permit direct comparison of changes in algal virus communities and their hosts.

To minimize the possibility that seasonal patterns were due to within-gel biases (as noted by Ferrari & Hollibaugh 1999), the order of samples loaded in the gradient gels was arbitrary. Despite limitations of fingerprint cluster analysis, gel comparison software was essential to this study; the normalization of individual samples and alignment of common bands within a fingerprint set was not possible without it. Computerized normalization and alignment permitted temporal ordering of fingerprints not run in successive gel lanes. The fingerprint approach described provided useful qualitative information on the temporal variability of algal viruses and eukaryotes; in all 3 fingerprint sets, bands appeared and disappeared in successive samples and, within each set, patterns and variability changed with time. In addition, the lack of coincidence between changes in algal virus and eukaryote community composition was demonstrated by cluster analysis of the fingerprints. The minimum similarity of any branches on AVS tree was higher than the minimum on the 18S trees and, when compared to 18S fingerprints, the AVS fingerprints clustered into groups with higher minimum similarity (Fig. 5). The cluster analyses also showed that the AVS fingerprints from the same season tended to cluster in larger groups than the 18S fingerprints.

There is accumulating evidence that some phytoplankton are infected by viruses not closely related to the Phycodnaviridae (Lawrence et al. 2001, Tai et al. 2002). In addition, our attempts to amplify pol genes from several viruses that are morphologically similar to the Phycodnaviridae have always failed. Thus, it is apparent that the viruses detected with the AVS primers used in this study likely represent a subset of the Phycodnaviridae. Therefore, the lack of coincidence between changes in algal virus and eukaryote communities at Jericho Pier was not surprising. Although they were not always coupled, changes in algal virus and eukaryote community composition were observed throughout the sample period, suggesting seasonality and succession in both communities. Nonetheless, through most of this study, eukaryote community composition was highly variable, yet algal virus community composition was relatively stable. This is the first study demonstrating that specific taxa of algal viruses are temporally stable and can persist in fluctuating physical and biological environments. This sug-
gests that the production of some taxa of algal-viruses is constant, and therefore they are a constant source of mortality to some phytoplankton. Hence, we speculate that these viruses influence phytoplankton community structure and may help maintain high host diversity.

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